Revised: January 2001
Prograf<sup>®</sup>
tacrolimus capsules
tacrolimus injection (for intravenous infusion only)

6

#### WARNING

Increased susceptibility to infection and the possible development of lymphoma may result from immunosuppression. Only physicians experienced in immunosuppressive therapy and management of organ transplant patients should prescribe Prograf. Patients receiving the drug should be managed in facilities equipped and staffed with adequate laboratory and supportive medical resources. The physician responsible for maintenance therapy should have complete information requisite for the follow-up of the patient.

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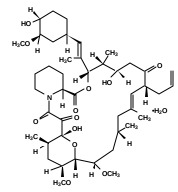
#### **DESCRIPTION:**

9 Prograf is available for oral administration as 10 capsules (tacrolimus capsules) containing the 11 equivalent of 0.5 mg, 1 mg or 5 mg of anhydrous 12 tacrolimus. Inactive ingredients include lactose, 13 hydroxypropyl methylcellulose, croscarmellose 14 sodium, and magnesium stearate. The 0.5 mg 15 capsule shell

5 capsule shell

16	contains gelatin, titanium dioxide and ferric oxide,
17	the 1 mg capsule shell contains gelatin and
18	titanium dioxide, and the 5 mg capsule shell
19	contains gelatin, titanium dioxide and ferric oxide.
20	
21	Prograf is also available as a sterile
22	solution (tacrolimus injection) containing the
23	equivalent of 5 mg anhydrous tacrolimus in 1 mL
24	for administration by intravenous infusion only.
25	Each mL contains polyoxyl 60 hydrogenated
26	castor oil (HCO-60), 200 mg, and dehydrated
27	alcohol, USP, 80.0% v/v. Prograf injection must
28	be diluted with 0.9% Sodium Chloride Injection
29	or 5% Dextrose Injection before use.
30	Tacrolimus, previously known as
31	FK506, is the active ingredient in Prograf.
32	Tacrolimus is a macrolide immunosuppressant
33	produced by Streptomyces tsukubaensis.
34	Chemically, tacrolimus is designated as [3S-
35	$[3R^*[E(1S^*,3S^*,4S^*)],4S^*,5R^*,8S^*,9E,12R^*,14R^*,$
36	15S*,16R*,18S*,19S*,26aR*]]-
37	5,6,8,11,12,13,14,15,16,17,18,19,24,25,26,26a
38	-hexadecahydro-5,19-dihydroxy-3-[2-(4-
39	hydroxy-3-methoxycyclohexyl)-1-
40	methylethenyl]-14,16-dimethoxy-4,10,12,18-
41	tetramethyl-8-(2-propenyl)-15,19-epoxy-3H-
42	pyrido[2,1-c][1,4] oxaazacyclotricosine-
43	1,7,20,21(4H,23H)-tetrone, monohydrate.
44	

The chemical structure of tacrolimus is:



Tacrolimus has an empirical formula of C<sub>44</sub>H<sub>69</sub>NO<sub>12</sub>CH<sub>2</sub>O and a formula weight of 822.05. Tacrolimus appears as white crystals or crystalline powder. It is practically insoluble in water, freely soluble in ethanol, and very soluble in methanol and chloroform.

### **CLINICAL PHARMACOLOGY:**

### Mechanism of Action

Tacrolimus prolongs the survival of the host and transplanted graft in animal transplant models of liver, kidney, heart, bone marrow, small bowel and pancreas, lung and trachea, skin, cornea, and limb.

In animals, tacrolimus has been demonstrated to suppress some humoral immunity and, to a greater extent, cell-mediated reactions such as allograft rejection, delayed type hypersensitivity, collagen- induced arthritis, experimental allergic encephalomyelitis, and graft

versus host disease.

78 79 Tacrolimus inhibits T-lymphocyte 80 activation, although the exact mechanism of action is not known. Experimental evidence suggests 81 that tacrolimus binds to an intracellular protein, 82 83 FKBP-12. A complex of tacrolimus-FKBP-12, 84 calcium, calmodulin, and calcineurin is then formed and the phosphatase activity of calcineurin 85 This effect may prevent the 86 inhibited. 87 dephosphorylation and translocation of nuclear 88 factor of activated T-cells (NF-AT), a nuclear 89 component thought to initiate gene transcription 90 for the formation of lymphokines (such as 91 interleukin-2, gamma interferon). The net result is the inhibition of T-lymphocyte activation (i.e., 92 93 immunosuppression). 94 95 **Pharmacokinetics** 96 Tacrolimus activity is primarily due to the parent 97 The pharmacokinetic parameters (mean"S.D.) of tacrolimus have been determined 98 99 following intravenous (IV) and oral (PO) 100 administration in healthy volunteers, and in kidney 101 transplant and liver transplant patients. (See table

102

below.)

103 104

Population	N	Route (Dose)	Parameters					
			C <sub>max</sub> (ng/mL)	T <sub>max</sub> (hr)	AUC (ng hr/mL)	t 2 (hr)	Cl (L/hr/kg	V (L/kg)
Healthy Volunteers	8	IV (0.025 mg/kg/4hr)	1		598* " 125	34.2 " 7.7	0.040 "0.009	1.91 "0.31
	16	PO (5 mg)	29.7 "7.2	1.6 "0.7	243** "73	34.8 "11.4	0.041 <b>H</b> "0.008	1.94 <b>H</b> "0.53
Kidney Transplant		IV (0.02 mg/kg/12hr)			294*** "262	18.8 "16.7	0.083 "0.050	1.41 "0.66
Pts	26	PO (0.2 mg/kg/day)	19.2 "10.3	3.0	203***	#	#	#
		PO (0.3 mg/kg/day)	24.2 "15.8	1.5	288*** "93	#	#	#
Liver Transplant Pts	17	IV (0.05 mg/kg/12 hr)	1		3300*** "2130	11.7 "3.9	0.053 "0.017	0.85 "0.30
		PO (0.3 mg/kg/day)	68.5 "30.0	2.3 "1.5	519*** "179	#	#	#

H Corrected for individual bioavailability

106 \* AUC<sub>0-120</sub> 107

\*\* AUC<sub>0-72</sub>

\*\*\* AUC<sub>0-inf</sub>

-- not applicable

# not available

110 111

108 109

105

Due to intersubject variability in tacrolimus 112

pharmacokinetics, individualization of dosing 113

regimen is necessary for optimal therapy. (See 114

DOSAGE AND ADMINISTRATION). 115

Pharmacokinetic data indicate that whole

117 blood concentrations rather than plasma 118 concentrations serve as the more appropriate 119 sampling compartment to describe tacrolimus 120 pharmacokinetics. 121 122 Absorption 123 Absorption of tacrolimus from the gastrointestinal 124 tract after oral administration is incomplete and 125 variable. The absolute bioavailability of tacrolimus was 17"10% in adult kidney 126 127 transplant patients (N=26), 22" 6% in adult liver 128 transplant patients (N=17), and 18"5% in 129 healthy volunteers (N=16). 130 A single dose study conducted in 32 131 healthy volunteers established the bioequivalence 132 of the 1 mg and 5 mg capsules. Another single 133 dose study in 32 healthy volunteers established 134 the bioequivalence of the 0.5 mg and 1 mg 135 capsules. Tacrolimus maximum blood 136 concentration (C<sub>max</sub>) and area under the curve 137 (AUC) appeared to increase in a dose-138 proportional fashion in 18 fasted healthy 139 volunteers receiving a single oral dose of 3, 7 and 10 mg. 140 141 In 18 kidney transplant patients, 142 tacrolimus trough concentrations from 3 to 30 ng/mL measured at 10-12 hours post-dose 143 144 (C<sub>min</sub>) correlated well with the AUC (correlation 145 coefficient 0.93). In 24 liver transplant patients 146 over a concentration range of 10 to 60 ng/mL, the correlation coefficient was 0.94. 147

148 149 Food Effects: The rate and extent of 150 tacrolimus absorption were greatest under fasted 151 conditions. The presence and composition of 152 food decreased both the rate and extent of 153 tacrolimus absorption when administered to 15 154 healthy volunteers. 155 The effect was most pronounced with a 156 high-fat meal (848 kcal, 46% fat): mean AUC 157 and C<sub>max</sub> were decreased 37% and 77%, 158 respectively; T<sub>max</sub> was lengthened 5-fold. A high-159 carbohydrate meal (668 kcal, 160 carbohydrate) decreased mean AUC and mean 161  $C_{\text{max}}$  by 28% and 65%, respectively. 162 In healthy volunteers (N=16), the time of 163 the meal also affected tacrolimus bioavailability. 164 When given immediately following the meal, 165 mean C<sub>max</sub> was reduced 71%, and mean AUC 166 was reduced 39%, relative to the fasted 167 condition. When administered 1.5 hours following the meal, mean C<sub>max</sub> was reduced 63%, 168 169 and mean AUC was reduced 39%, relative to the 170 fasted condition. 171 In 11 liver transplant patients, Prograf 172 administered 15 minutes after a high fat (400 173 kcal, 34% fat) breakfast, resulted in decreased AUC (27" 18%) and  $C_{max}$  (50"19%), as 174

compared to a fasted state.

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176	
177	
178	Distribution
179	The plasma protein binding of tacrolimus is
180	approximately 99% and is independent of
181	concentration over a range of 5-50 ng/mL.
182	Tacrolimus is bound mainly to albumin and alpha-
183	1-acid glycoprotein, and has a high level of
184	association with erythrocytes. The distribution of
185	tacrolimus between whole blood and plasma
186	depends on several factors, such as hematocrit,
187	temperature at the time of plasma separation,
188	drug concentration, and plasma protein
189	concentration. In a U.S. study, the ratio of whole
190	blood concentration to plasma concentration
191	averaged 35 (range 12 to 67).
192	
193	<u>Metabolism</u>
194	Tacrolimus is extensively metabolized by the
195	mixed-function oxidase system, primarily the
196	cytochrome P-450 system (CYP3A). A
197	metabolic pathway leading to the formation of 8
198	possible metabolites has been proposed.
199	Demethylation and hydroxylation were identified
200	as the primary mechanisms of biotransformation
201	in vitro. The major metabolite identified in
202	incubations with human liver microsomes is 13-
203	demethyl tacrolimus. In in vitro studies, a 31-
204	demethyl metabolite has been reported to have
205	the same activity as tacrolimus.

206 207 208 Excretion 209 The mean clearance following IV administration 210 of tacrolimus is 0.040, 0.083 and 0.053 L/hr/kg 211 in healthy volunteers, adult kidney transplant 212 patients and adult liver transplant patients, 213 respectively. In man, less than 1% of the dose 214 administered is excreted unchanged in urine. 215 In a mass balance study of IV administered radiolabeled tacrolimus to 6 healthy 216 217 volunteers, the mean recovery of radiolabel was 77.8" 12.7%. Fecal elimination accounted for 218 219 92.4" 1.0% and the elimination half-life based on 220 radioactivity was 48.1" 15.9 hours whereas it 221 was 43.5"11.6 hours based on tacrolimus 222 concentrations. The mean clearance of radiolabel 223 was 0.029" 0.015 L/hr/kg and clearance of tacrolimus was 0.029" 0.009 L/hr/kg. When 224 225 administered PO, the mean recovery of the 226 radiolabel was 94.9" 30.7%. Fecal elimination 227 accounted for 92.6" 30.7%, urinary elimination accounted for 2.3" 1.1% and the elimination half-228 229 life based on radioactivity was 31.9" 10.5 hours whereas it was 48.4" 12.3 hours based on 230 231 tacrolimus concentrations. The mean clearance 232 of radiolabel was 0.226" 0.116 L/hr/kg and

clearance of tacrolimus 0.172" 0.088 L/hr/kg.

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234	
235	Special Populations
236	<u>Pediatric</u>
237	Pharmacokinetics of tacrolimus have been studied
238	in liver transplantation patients, 0.7 to 13.2 years
239	of age. Following IV administration of a 0.037
240	mg/kg/day dose to 12 pediatric patients, mean
241	terminal half-life, volume of distribution and
242	clearance were 11.5"3.8 hours, 2.6"2.1 L/kg
243	and 0.138" 0.071 L/hr/kg, respectively.
244	Following oral administration to 9 patients, mean
245	AUC and C <sub>max</sub> were 337" 167 ng\$hr/mL and
246	43.4" 27.9 ng/mL, respectively. The absolute
247	bioavailability was 31" 21%.
248	Whole blood trough concentrations from
249	31 patients less than 12 years old showed that
250	pediatric patients needed higher doses than adults
251	to achieve similar tacrolimus trough
252	concentrations. (See DOSAGE AND
253	ADMINISTRATION).
254	
255	Renal and Hepatic Insufficiency
256	The mean pharmacokinetic parameters for
257	tacrolimus following single administrations to
258	patients with renal and hepatic impairment are
259	given in the following table.

260

Population	Dose	AUC <sub>0-t</sub>	t <sub>1/2</sub>	V	Cl
(No. of Patients)		(ng • hr/mL)	(hr)	(L/kg)	(L/hr/kg)
Renal	0.02				
Impairment	mg/kg/4hr	393±123	26.3±9.2	1.07	0.038
(n=12)	IV	(t=60 hr)		±0.20	±0.014
Mild Hepatic	0.02	367±107	60.6±43.8	3.1	0.042
Impairment	mg/kg/4hr	(t=72 hr)	Range: 27.8 – 141	±1.6	±0.02
(n=6)	IV				
	7.7 mg	488±320	66.1±44.8	3.7	0.034
	PO	(t=72 hr)	Range: 29.5 – 138	±4.7*	±0.019*
Severe	0.02 mg/kg/4hr	762±204			
Hepatic	IV (n=2)	(t=120 hr)	198±158		
Impairment			Range: 81-436		
(n=6, IV)	0.01 mg/kg/8hr	289±117	-	3.9±1.0	0.017±0.013
	IV (n=4)	(t=144 hr)			
(n=5, PO)†	8 mg PO	658			
	(n=1)	(t=120 hr)	119±35		
			Range: 85-178	3.1±3.4*	0.016±0.011*
	5 mg PO	533±156			
	(n=4)	(t=144 hr)			
	4 mg PO				
	(n=1)				

\* corrected for bioavailability

† 1 patient did not receive the PO dose

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## Renal Insufficiency:

- 265 Tacrolimus pharmacokinetics following a single
- 266 IV administration were determined in 12 patients
- 267 (7 not on dialysis and 5 on dialysis, serum
- 268 creatinine of 3.9"1.6 and 12.0"2.4 mg/dL,
- 269 respectively) prior to their kidney transplant. The
- 270 pharmacokinetic parameters obtained were
- similar for both groups.

272	
273	The mean clearance of tacrolimus in
274	patients with renal dysfunction was similar to that
275	in normal volunteers (see previous table).
276	
277	Hepatic Insufficiency:
278	Tacrolimus pharmacokinetics have been
279	determined in six patients with mild hepatic
280	dysfunction (mean Pugh score: 6.2) following
281	single IV and oral administrations. The mean
282	clearance of tacrolimus in patients with mild
283	hepatic dysfunction was not substantially different
284	from that in normal volunteers (see previous
285	table). Tacrolimus pharmacokinetics were
286	studied in 6 patients with severe hepatic
287	dysfunction (mean Pugh score:>10). The mean
288	clearance was substantially lower in patients with
289	severe hepatic dysfunction, irrespective of the
290	route of administration.
291	
292	Race
293	A formal study to evaluate the pharmacokinetic
294	disposition of tacrolimus in Black transplant
295	patients has not been conducted. However, a
296	retrospective comparison of Black and Caucasian
297	kidney transplant patients indicated that Black
298	patients required higher tacrolimus doses to attain
299	similar trough concentrations. (See DOSAGE
300	AND ADMINISTRATION)

301	
302	
303	Gender
304	A formal study to evaluate the effect of gender or
305	tacrolimus pharmacokinetics has not been
306	conducted, however, there was no difference in
307	dosing by gender in the kidney transplant trial. A
308	retrospective comparison of pharmacokinetics in
309	healthy volunteers, and in kidney and liver
310	transplant patients indicated no gender-based
311	differences.
312	
313	<u>Clinical Studies</u>
314	Liver Transplantation
315	The safety and efficacy of Prograf-based
316	immunosuppression following orthotopic liver
317	transplantation were assessed in two prospective,
318	randomized, non-blinded multicenter studies. The
319	active control groups were treated with a
320	cyclosporine-based immunosuppressive regimen.
321	Both studies used concomitant adrenal
322	corticosteroids as part of the immunosuppressive
323	regimens. These studies were designed to
324	evaluate whether the two regimens were
325	therapeutically equivalent, with patient and graft
326	survival at 12 months following transplantation as
327	the primary endpoints. The Prograf-based
328	immunosuppressive regimen was found to be
329	equivalent to the cyclosporine-based
330	immunosuppressive regimens.

In one trial, 529 patients were enrolled at 12 clinical sites in the United States; prior to surgery, 263 were randomized to the Prograf-based immunosuppressive regimen and 266 to a cyclosporine-based immunosuppressive regimen (CBIR). In 10 of the 12 sites, the same CBIR protocol was used, while 2 sites used different control protocols. This trial excluded patients with renal dysfunction, fulminant hepatic failure with Stage IV encephalopathy, and cancers; pediatric patients (< 12 years old) were allowed. 

In the second trial, 545 patients were enrolled at 8 clinical sites in Europe; prior to surgery, 270 were randomized to the Prograf-based immunosuppressive regimen and 275 to CBIR. In this study, each center used its local standard CBIR protocol in the active-control arm. This trial excluded pediatric patients, but did allow enrollment of subjects with renal dysfunction, fulminant hepatic failure in Stage IV encephalopathy, and cancers other than primary hepatic with metastases.

One-year patient survival and graft survival in the Prograf-based treatment groups were equivalent to those in the CBIR treatment groups in both studies. The overall one-year patient survival (CBIR and Prograf-based treatment groups combined) was 88% in the U.S. study and 78% in the European study.

361 The overall one-year graft survival (CBIR and 362 Prograf-based treatment groups combined) was 81% in the U.S. study and 73% in the European 363 364 study. In both studies, the median time to convert 365 from IV to oral Prograf dosing was 2 days. Because of the nature of the study design, 366 367 comparisons of differences in secondary 368 endpoints, such as incidence of acute rejection, 369 refractory rejection or use of OKT3 for steroid-370 resistant rejection, could not be reliably made. 371 372 Kidney Transplantation 373 Prograf-based immunosuppression following 374 kidney transplantation was assessed in a Phase randomized. 375 multicenter, non-blinded, 376 prospective study. There were 412 kidney transplant patients enrolled at 19 clinical sites in 377 378 the United States. Study therapy was initiated 379 when renal function was stable as indicated by a 380 serum creatinine < 4 mg/dL (median of 4 days 381 after transplantation, range 1 to 14 days). 382 Patients less than 6 years of age were excluded. 383 There were 205 patients randomized to Prograf-based immunosuppression and 207 384 385 patients were randomized to cyclosporine-based 386 immunosuppression. All patients received prophylactic induction therapy consisting of an 387 388 antilymphocyte antibody preparation,

corticosteroids and azathioprine.

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Overall one year patient and graft survival was

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391	96.1% and 89.6%, respectively and was
392	equivalent between treatment arms.
393	Because of the nature of the study design,
394	comparisons of differences in secondary
395	endpoints, such as incidence of acute rejection,
396	refractory rejection or use of OKT3 for steroid-
397	resistant rejection, could not be reliably made.
398	
399	INDICATIONS AND USAGE:
400	Prograf is indicated for the prophylaxis of organ
401	rejection in patients receiving allogeneic liver or
402	kidney transplants. It is recommended that
403	Prograf be used concomitantly with adrenal
404	corticosteroids. Because of the risk of
405	anaphylaxis, Prograf injection should be reserved
406	for patients unable to take Prograf capsules
407	orally.
408	
409	CONTRAINDICATIONS:
410	Prograf is contraindicated in patients with a
411	hypersensitivity to tacrolimus. Prograf injection is
412	contraindicated in patients with a hypersensitivity
413	to HCO-60 (polyoxyl 60 hydrogenated castor
414	oil).
415	
416	WARNINGS:
417	(See boxed <b>WARNING.</b> )
418	Insulin-dependent post-transplant diabetes
419	mellitus (PTDM) was reported in 20% of
420	Prograf-treated kidney transplant patients

421	without pretransplant history of diabetes mellitus
422	in the Phase III study (See Tables Below). The
423	median time to onset of PTDM was 68 days.
424	Insulin dependence was reversible in 15% of
425	these PTDM patients at one year and in 50% at
426	two years post transplant. Black and Hispanic
427	kidney transplant patients were at an increased
428	risk of development of PTDM.

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# Incidence of Post Transplant Diabetes Mellitus and Insulin Use at 2 Years in Kidney Transplant Recipients in the Phase

III Study

Status of PTDM*	Prograf	CBIR
Patients without pretransplant history of diabetes mellitus.	151	151
New onset PTDM*, 1st Year	30/151 (20%)	6/151 (4%)
Still insulin dependent at one year in those without prior history of diabetes.	25/151(17%)	5/151 (3%)
New onset PTDM* post 1 year	1	0
Patients with PTDM* at 2 years	16/151 (11%)	5/151 (3%)

\*use of insulin for 30 or more consecutive days, with <

435 5 day gap, without a prior history of insulin dependent

436 diabetes mellitus or non insulin dependent diabetes

437 mellitus.

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**Development of Post Transplant Diabetes** 

Mellitus by Race and by Treatment Group

during First Year Post Kidney

Transplantation in the Phase III Study

Patient	Prograf		CBIR	
Race	No. of Patients at Risk	Patients Who Developed PTDM*	No. of Patients At Risk	Patients Who Developed PTDM*
Black	41	15 (37%)	36	3 (8%)
Hispanic	17	5 (29%)	18	1 (6%)
Caucasian	82	10 (12%)	87	1 (1%)
Other	11	0 (0%)	10	1 (10%)
Total	151	30 (20%)	151	6 (4%)

st use of insulin for 30 or more consecutive days, with

445 < 5 day gap, without a prior history of insulin

dependent diabetes mellitus or non insulin dependent

447 diabetes mellitus.

448	Insulin-dependent post-transplant diabetes
449	mellitus was reported in 18% and 11% of
450	Prograf-treated liver transplant patients and
451	was reversible in 45% and 31% of these
452	patients at one year post transplant, in the
453	U.S. and European randomized studies,
454	respectively (See Table below).
455	Hyperglycemia was associated with the use of
456	Prograf in 47% and 33% of liver transplant
457	recipients in the U.S. and European randomized
458	studies, respectively, and may require treatment
459	(see ADVERSE REACTIONS).

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## **Incidence of Post Transplant Diabetes** Mellitus and Insulin Use at One Year in **Liver Transplant Recipients**

Status of PTDM*	US Study		European Study		
	Prograf	CBIR	Prograf	CBIR	
Patients at risk **	239	236	239	249	
New Onset PTDM*	42 (18%)	30 (13%)	26 (11%)	12(5%)	
Patients still on insulin at 1 year	23 (10%)	19 (8%)	18 (8%)	6 (2%)	

464 465

\* use of insulin for 30 or more consecutive days,

with < 5 day gap, without a prior history of

insulin dependent diabetes mellitus or non

467 insulin dependent diabetes mellitus.

468 \*\*Patients without pretransplant history of diabetes mellitus.

469 470

466

471 **Prograf** can cause neurotoxicity and 472 nephrotoxicity, particularly when used in high 473 Nephrotoxicity was reported in doses. 474 approximately 52% of kidney transplantation 475 patients and in 40% and 36% of liver transplantation patients receiving Prograf in the 476 477 European randomized U.S. and trials, 478 respectively (see ADVERSE REACTIONS). 479 More overt nephrotoxicity is seen early after 480 transplantation, characterized by increasing serum creatinine and a decrease in urine output. 481 482 Patients with impaired renal function should be 483 monitored closely as the dosage of Prograf may 484 need to be reduced. In patients with persistent 485 elevations of serum creatinine who are 486 unresponsive to dosage adjustments, 487 consideration should be given to changing to 488 another immunosuppressive therapy. 489 should be taken in using tacrolimus with other 490 nephrotoxic drugs. In particular, to avoid 491 excess nephrotoxicity, Prograf should not be 492 used simultaneously with cyclosporine. 493 Prograf or cyclosporine should discontinued at least 24 hours prior to 494 495 initiating the other. In the presence of 496 elevated **Prograf** or cyclosporine 497 concentrations, dosing with the other drug 498 usually should be further delayed.

499 500 Mild to severe hyperkalemia was 501 reported in 31% of kidney transplant recipients and in 45% and 13% of liver transplant recipients 502 treated with Prograf in the U.S. and European 503 randomized trials, respectively, and may require 504 505 treatment (see ADVERSE REACTIONS). 506 Serum potassium levels should be monitored 507 and potassium-sparing diuretics should not 508 be used during Prograf therapy (see 509 PRECAUTIONS). 510 Neurotoxicity, including tremor, 511 headache, and other changes in motor function, 512 mental status, and sensory function were reported in approximately 55% of liver transplant 513 514 recipients in the two randomized studies. Tremor 515 occurred more often in Prograf-treated kidney 516 transplant patients (54%) compared 517 cyclosporine-treated patients. The incidence of other neurological events in kidney transplant 518 519 patients was similar in the two treatment groups 520 (see ADVERSE REACTIONS). Tremor and 521 headache have been associated with high whole-522 blood concentrations of tacrolimus and may 523 respond to dosage adjustment. Seizures have 524 occurred in adult and pediatric patients receiving 525 Prograf (see ADVERSE REACTIONS). 526 Coma and delirium also have been associated 527 with high plasma concentrations of tacrolimus. 528

529	As in patients receiving other
530	immunosuppressants, patients receiving Prograf
531	are at increased risk of developing lymphomas
532	and other malignancies, particularly of the skin.
533	The risk appears to be related to the intensity and
534	duration of immunosuppression rather than to the
535	use of any specific agent. A lymphoproliferative
536	disorder (LPD) related to Epstein-Barr Virus
537	(EBV) infection has been reported in
538	immunosuppressed organ transplant recipients.
539	The risk of LPD appears greatest in young
540	children who are at risk for primary EBV
541	infection while immunosuppressed or who are
542	switched to Prograf following long-term
543	immunosuppression therapy. Because of the
544	danger of oversuppression of the immune system
545	which can increase susceptibility to infection,
546	combination immunosuppressant therapy should
547	be used with caution.
548	A few patients receiving Prograf injection
549	have experienced anaphylactic reactions.
550	Although the exact cause of these reactions is not
551	known, other drugs with castor oil derivatives in
552	the formulation have been associated with
553	anaphylaxis in a small percentage of patients.
554	Because of this potential risk of anaphylaxis,
555	Prograf injection should be reserved for patients
556	who are unable to take Prograf capsules.

557

558 **Patients receiving Prograf injection** 559 should be under continuous observation for 560 at least the first 30 minutes following the 561 start of the infusion and at frequent intervals If signs or symptoms of 562 thereafter. 563 anaphylaxis occur, the infusion should be 564 stopped. An aqueous solution of epinephrine 565 should be available at the bedside as well as 566 a source of oxygen. 567 568 569 **PRECAUTIONS:** 570 General 571 Hypertension is a common adverse effect of 572 **Prograf** therapy (see **ADVERSE** 573 **REACTIONS**). Mild or moderate hypertension 574 is more frequently reported than severe 575 hypertension. Antihypertensive therapy may be required; the control of blood pressure can be 576 577 accomplished with any of the common 578 antihypertensive agents. Since tacrolimus may 579 cause hyperkalemia, potassium-sparing diuretics should be avoided. While calcium-channel 580 581 blocking agents can be effective in treating 582 Prograf-associated hypertension, care should be taken since interference with tacrolimus 583 584 metabolism may require a dosage reduction (see 585 Drug Interactions). 586

## 587 Renally and Hepatically Impaired Patients 588 For patients with renal insufficiency some 589 evidence suggests that lower doses should be

590 used (see CLINICAL PHARMACOLOGY

and DOSAGE AND ADMINISTRATION).
 The use of Prograf in liver transplant

The use of Prograf in liver transplant recipients experiencing post-transplant hepatic impairment may be associated with increased risk of developing renal insufficiency related to high whole-blood levels of tacrolimus. These patients should be monitored closely and dosage adjustments should be considered. Some evidence suggests that lower doses should be used in these patients (see **DOSAGE AND ADMINISTRATION**).

### Myocardial Hypertrophy

Myocardial hypertrophy has been reported in association with the administration of Prograf, and is generally manifested by echocardiographically demonstrated concentric increases in left ventricular posterior wall and interventricular septum thickness. Hypertrophy has been observed in infants, children and adults. This condition appears reversible in most cases following dose reduction or discontinuance of therapy. In a group of 20 patients with pre- and post-treatment echocardiograms who showed evidence of myocardial hypertrophy, mean tacrolimus

616 tacrolimus

whole blood concentrations during the period prior to diagnosis of myocardial hypertrophy ranged from 11 to 53 ng/mL in infants (N=10, age 0.4 to 2 years), 4 to 46 ng/mL in children (N=7, age 2 to 15 years) and 11 to 24 ng/mL in adults (N=3, age 37 to 53 years).

In patients who develop renal failure or clinical manifestations of ventricular dysfunction while receiving Prograf therapy, echocardiographic evaluation should be considered. If myocardial hypertrophy is diagnosed, dosage reduction or discontinuation of Prograf should be considered.

### Information for Patients

Patients should be informed of the need for repeated appropriate laboratory tests while they are receiving Prograf. They should be given complete dosage instructions, advised of the potential risks during pregnancy, and informed of the increased risk of neoplasia. Patients should be informed that changes in dosage should not be undertaken without first consulting their physician.

Patients should be informed that Prograf can cause diabetes mellitus and should be advised of the need to see their physician if they develop frequent urination, increased thirst or hunger.

645	Laboratory Tests
646	Serum creatinine, potassium, and fasting glucose
647	should be assessed regularly. Routine monitoring
648	of metabolic and hematologic systems should be
649	performed as clinically warranted.
650	
651	Drug Interactions
652	Due to the potential for additive or synergistic
653	impairment of renal function, care should be taken
654	when administering Prograf with drugs that may
655	be associated with renal dysfunction. These
656	include, but are not limited to, aminoglycosides,
657	amphotericin B, and cisplatin. Initial clinica
658	experience with the co-administration of Progra
659	and cyclosporine resulted in additive/synergistic
660	nephrotoxicity. Patients switched from
661	cyclosporine to Prograf should receive the first
662	Prograf dose no sooner than 24 hours after the
663	last cyclosporine dose. Dosing may be further
664	delayed in the presence of elevated cyclosporine
665	levels.
666	
667	Drugs that May Alter Tacrolimus
668	Concentrations
669	Since tacrolimus is metabolized mainly by the
670	CYP3A enzyme systems, substances known to
671	inhibit these enzymes may decrease the
672	metabolism or increase bioavailability of
673	tacrolimus as indicated by increased whole blood
674	or plasma concentrations. Drugs known

to induce these enzyme systems may result in an increased metabolism of tacrolimus or decreased bioavailability as indicated by decreased whole blood or plasma concentrations. Monitoring of blood concentrations and appropriate dosage adjustments are essential when such drugs are used concomitantly.

## \*Drugs That May Increase Tacrolimus Blood Concentrations:

Calcium	Antifungal	Macrolide
Channel Blockers	Agents	<b>Antibiotics</b>
diltiazem	clotrimazole	clarithromycin
nicardipine	fluconazole	erythromycin
nifedipine	itraconazole	troleandomycin
verapamil	ketoconazole	

Gastrointestinal	Other
Prokinetic	Drugs
Agents	bromocriptine
cisapride	cimetidine
metoclopramide	cyclosporine
	danazol
	ethinyl estradiol
	methylprednisolone
	omeprazole
	protease inhibitors
	nefazodone

In a study of 6 normal volunteers, a significant increase in tacrolimus oral bioavailability ( $14\pm5\%$  vs.  $30\pm8\%$ ) was observed with concomitant ketoconazole administration (200 mg). The apparent oral clearance of tacrolimus during ketoconazole administration was significantly decreased

compared to tacrolimus alone (0.430±0.129 L/hr/kg vs. 0.148±0.043 L/hr/kg). Overall, IV clearance of tacrolimus was not significantly changed by ketoconazole co-administration, although it was highly variable between patients.

#### \*Drugs That May Decrease Tacrolimus Blood Concentrations:

 Anticonvulsants
 Antibiotics

 carbamazepine
 rifabutin

 phenobarbital
 rifampin

 phenytoin
 rifampin

#### **Herbal Preparations**

St. John's Wort

\*This table is not all inclusive.

St. John's Wort (hypericum perforatum) induces CYP3A4 and P-glycoprotein. Since tacrolimus is a substrate for CYP3A4, there is the potential that the use of St. John's Wort in patients receiving Prograf could result in reduced tacrolimus levels.

In a study of 6 normal volunteers, a significant decrease in tacrolimus oral bioavailability (14±6% vs. 7±3%) was observed with concomitant rifampin administration (600 mg). In addition, there was a significant increase in tacrolimus clearance (0.036±0.008 L/hr/kg vs. 0.053±0.010 L/hr/kg) with concomitant rifampin administration.

743	Interaction studies with drugs used in
744	HIV therapy have not been conducted.
745	However, care should be exercised when drugs
746	that are nephrotoxic (e.g., ganciclovir) or that are
747	metabolized by CYP3A (e.g., ritonavir) are
748	administered concomitantly with tacrolimus.
749	Tacrolimus may affect the pharmacokinetics of
750	other drugs (e.g., phenytoin) and increase their
751	concentration. Grapefruit juice affects CYP3A-
752	mediated metabolism and should be avoided
753	(See DOSAGE AND ADMINISTRATION).
754	
755	Other Drug Interactions
756	Immunosuppressants may affect vaccination.
757	Therefore, during treatment with Prograf,
758	vaccination may be less effective. The use of live
759	vaccines should be avoided; live vaccines may
760	include, but are not limited to measles, mumps,
761	rubella, oral polio, BCG, yellow fever, and TY
762	21a typhoid. <sup>1</sup>
763	
764	Carcinogenesis, Mutagenesis and
765	Impairment of Fertility
766	An increased incidence of malignancy is a
767	recognized complication of immunosuppression in
768	recipients of organ transplants. The most
769	common forms of neoplasms are non-Hodgkin's
770	lymphomas and carcinomas of the skin. As with
771	other immunosuppressive therapies, the risk of
772	malignancies in Prograf recipients may be higher
773	than in the normal, healthy

population. Lymphoproliferative disorders associated with Epstein-Barr Virus infection have been seen. It has been reported that reduction or discontinuation of immunosuppression may cause the lesions to regress.

No evidence of genotoxicity was seen in bacterial (*Salmonella* and *E. coli*) or mammalian (Chinese hamster lung-derived cells) in vitro assays of mutagenicity, the in vitro CHO/HGPRT assay of mutagenicity, or in vivo clastogenicity assays performed in mice; tacrolimus did not cause unscheduled DNA synthesis in rodent hepatocytes.

Carcinogenicity studies were carried out in male and female rats and mice. In the 80-week mouse study and in the 104-week rat study no relationship of tumor incidence to tacrolimus dosage was found. The highest doses used in the mouse and rat studies were 0.8 - 2.5 times (mice) and 3.5 - 7.1 times (rats) the recommended clinical dose range of 0.1 - 0.2 mg/kg/day when corrected for body surface area.

No impairment of fertility was demonstrated in studies of male and female rats. Tacrolimus, given orally at 1.0 mg/kg

799 (0.7 - 1.4X the recommended clinical dose 800 range of 0.1 - 0.2 mg/kg/day based on body 801 surface area corrections) to male and female rats, 802 prior to and during mating, as well as to dams during gestation and lactation, was associated 803 with embryolethality and with adverse effects on 804 805 female reproduction. Effects on female 806 reproductive function (parturition) 807 embryolethal effects were indicated by a higher 808 rate of pre-implantation loss and increased 809 numbers of undelivered and nonviable pups. 810 When given at 3.2 mg/kg (2.3 - 4.6X the 811 recommended clinical dose range based on body 812 surface area correction), tacrolimus was 813 associated with maternal and paternal toxicity as 814 well as reproductive toxicity including marked 815 adverse effects on estrus cycles, parturition, pup 816 viability, and pup malformations.

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### Pregnancy: Category C

In reproduction studies in rats and rabbits, adverse effects on the fetus were observed mainly at dose levels that were toxic to dams. Tacrolimus at oral doses of 0.32 and 1.0 mg/kg during organogenesis in rabbits was associated with maternal toxicity as well as an increase in incidence of abortions; these doses are equivalent to 0.5 - 1X and 1.6 - 3.3X the recommended clinical dose range (0.1 - 0.2 mg/kg) based on body surface area

829	corrections. At the higher dose only, an
830	increased incidence of malformations and
831	developmental variations was also seen.
832	Tacrolimus, at oral doses of 3.2 mg/kg during
833	organogenesis in rats, was associated with
834	maternal toxicity and caused an increase in late
835	resorptions, decreased numbers of live births, and
836	decreased pup weight and viability. Tacrolimus,
837	given orally at 1.0 and 3.2 mg/kg (equivalent to
838	0.7 - 1.4X and 2.3 - 4.6X the recommended
839	clinical dose range based on body surface area
840	corrections) to pregnant rats after organogenesis
841	and during lactation, was associated with reduced
842	pup weights.
843	No reduction in male or female fertility
844	was evident.
845	There are no adequate and well-
846	controlled studies in pregnant women.
847	Tacrolimus is transferred across the placenta.
848	The use of tacrolimus during pregnancy has been
849	associated with neonatal hyperkalemia and renal
850	dysfunction. Prograf should be used during
851	pregnancy only if the potential benefit to the
852	mother justifies potential risk to the fetus.
853	
854	Nursing Mothers
855	Since tacrolimus is excreted in human milk,
856	nursing should be avoided.

857	
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859	Pediatric Patients
860	Experience with Prograf in pediatric kidney
861	transplant patients is limited. Successful liver
862	transplants have been performed in pediatric
863	patients (ages up to 16 years) using Prograf. Two
864	randomized active-controlled trials of Prograf in
865	primary liver transplantation included 56
866	pediatric patients. Thirty-one patients were
867	randomized to Prograf-based and 25 to
868	cyclosporine-based therapies. Additionally, a
869	minimum of 122 pediatric patients were studied in
870	an uncontrolled trial of tacrolimus in living related
871	donor liver transplantation. Pediatric patients
872	generally required higher doses of Prograf to
873	maintain blood trough concentrations of
874	tacrolimus similar to adult patients (see
875	DOSAGE AND ADMINISTRATION).
876	
877	ADVERSE REACTIONS:
878	Liver Transplantation
879	The principal adverse reactions of Prograf are
880	tremor, headache, diarrhea, hypertension, nausea
881	and renal dysfunction. These occur with oral and
882	IV administration of Prograf and may respond to
883	a reduction in dosing. Diarrhea was sometimes
884	associated with other gastrointestinal complaints
885	such as nausea and vomiting.

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Hyperkalemia and hypomagnesemia have occurred in patients receiving Prograf therapy. Hyperglycemia has been noted in many patients; some may require insulin therapy (see **WARNINGS**).

The incidence of adverse events was determined in two randomized comparative liver transplant trials among 514 patients receiving tacrolimus and steroids and 515 patients receiving a cyclosporine-based regimen (CBIR). The proportion of patients reporting more than one adverse event was 99.8% in the tacrolimus group and 99.6% in the CBIR group. Precautions must be taken when comparing the incidence of adverse events in the U.S. study to that in the European study. The 12-month posttransplant information from the U.S. study and from the European study is presented below. The two studies also included different patient populations and patients were treated with immunosuppressive regimens of differing intensities. Adverse events reported in \$15% in tacrolimus patients (combined study results) are presented below for the two controlled trials in liver transplantation:

912					
913					
914	LIVER TRANSPLANTATION: ADVE	RSE			
915	EVENTS OCCURRING IN \$ 15% O	F			
916	PROGRAF-TREATED PATIENTS	_			
917	TROOKAT-TREATED TATEL(1)				
918 919 920		U.S. STUI			AN STUDY (%)
920		Prograf (N=250)	CBIR (N=250)	Prograf (N=264)	CBIR (N=265)
921		(11-230)	(11-230)	(11-204)	(11-203)
922	Nervous System				
923	Headache (See WARNINGS)	64	60	37	26
924	Tremor (See WARNINGS)	56	46	48	32
925	Insomnia	64	68	32	23
926	Paresthesia	40	30	17	17
927					
928	<u>Gastrointestinal</u>				
929	Diarrhea	72	47	37	27
930	Nausea	46	37	32	27
931	Constipation	24	27	23	21
932	LFT Abnormal	36	30	6	5
933	Anorexia	34	24	7	5
934	Vomiting	27	15	14	11
935					
936	<u>Cardiovascular</u>				
937	Hypertension (See PRECAUTIONS)	47	56	38	43
938					
939	<u>Urogenital</u>				
940	Kidney Function Abnormal (See WARNINGS)	40	27	36	23
941	Creatinine Increased (See WARNINGS)	39	25	24	19
942	BUN Increased (See WARNINGS)	30	22	12	9
943	Urinary Tract Infection	16	18	21	19
944	Oliguria	18	15	19	12
945					
946	Metabolic and Nutritional				
947	Hyperkalemia (See WARNINGS)	45	26	13	9
948	Hypokalemia	29	34	13	16
949	Hyperglycemia (See WARNINGS)	47	38	33	22
950	Hypomagnesemia	48	45	16	9

951					
952					
953					
954	Hemic and Lymphatic				
955	Anemia	47	38	5	1
956	Leukocytosis	32	26	8	8
957	Thrombocytopenia	24	20	14	19
958					
959	<u>Miscellaneous</u>				
960	Abdominal Pain	59	54	29	22
961	Pain	63	57	24	22
962	Fever	48	56	19	22
963	Asthenia	52	48	11	7
964	Back Pain	30	29	17	17
965	Ascites	27	22	7	8
966	Peripheral Edema	26	26	12	14
967					
968	Respiratory System				
969	Pleural Effusion	30	32	36	35
970	Atelectasis	28	30	5	4
971	Dyspnea	9	23	5	4
972					
973	Skin and Appendages				_
974	Pruritus	36	20	15	7
975	Rash	24	19	10	4
976					
977	Less frequently observed adverse reaction	ns in			
978	both liver transplantation and kidney				
979	transplantation patient are described unde	r the			
980	subsection Less Frequently Reported				
981	Adverse Reactions below.				
	Auverse Reactions below.				
982					
983	Kidney Transplantation				
984	The most common adverse reactions repo	orted			
985	were infection, tremor, hypertension, dec				
	* <del>-</del>				
986	renal function, constipation, diarrhea, hea	uacne,			
987	abdominal pain and insomnia.				

988			
989	Adverse events th	at occurred in \$ 15	
990	% of Prograf-treated kidn	ey transplant patients	
991	are presented below:		
992	1		
993	KIDNEY		
994	TRANSPLANTATION:		
995	ADVERSE EVENTS		
996	OCCURRING IN \$ 15%		
997	OF PROGRAF-		
998	TREATED PATIENTS		
1000		<b>D</b> (	CDID
1001 1002		Prograf ( <u>N=205</u> )	CBIR (N=207)
1002	Nervous System	<u>(11–203)</u>	(11-201)
1003	Tremor (See		
1004	WARNINGS)	54	34
1006	Headache (See		
1007	WARNINGS)	44	38
1008	Insomnia	32	30
1009	Paresthesia	23	16
1010	Dizziness	19	16
1011			
1012	<u>Gastrointestinal</u>		
1013	Diarrhea	44	41
1014	Nausea	38	36
1015	Constipation	35	43
1016	Vomiting	29	23
1017	Dyspepsia	28	20
1018 1019	Cardiovascular		
1020	Hypertension (See		
1020	PRECAUTIONS)	50	52
1022	Chest pain	19	13

1023			
1024	<u>Urogenital</u>		
1025	Creatinine increased		
1026	(See WARNINGS)	45	42
1027	Urinary tract infection	34	35
1028			
1029	Metabolic and Nutritional		
1030	Hypophosphatemia	49	53
1031	Hypomagnesemia	34	17
1032	Hyperlipemia	31	38
1033	Hyperkalemia (See		
1034	WARNINGS)	31	32
1035	Diabetes mellitus		
1036	(See WARNINGS)	24	9
1037	Hypokalemia	22	25
1038	Hyperglycemia (See	22	1.6
1039	WARNINGS)	22	16
1040	Edema	18	19
1041			
1042	Hemic and Lymphatic		
1043	Anemia	30	24
1044	Leukopenia	15	17
1045			
1046	<u>Miscellaneous</u>		
1047	Infection	45	49
1048	Peripheral edema	36	48
1049	Asthenia	34	30
1050	Abdominal pain	33	31
1051	Pain	32	30
1052	Fever	29	29
1053	Back pain	24	20

1054 1055			
1056	Respiratory System		
1057	Dyspnea	22	18
1058	Cough increased	18	15
1059	-		
1060	<u>Musculoskeletal</u>		
1061	Arthralgia	25	24
1062			
1063	<u>Skin</u>		
1064 1065	Rash Pruritis	17	12 7
1065	PTUTIUS	15	/
1067	Less frequently observed	ved adverse r	eactions in
1068	both liver transp	plantation ar	d kidney
1069	transplantation patients	s are described	l under the
1070	subsection Less I	Frequently	Reported
1071	Adverse Reactions s	hown below.	
10/1	ria verbe reactions s	110 11111 0010 111.	
1072	riuverse reactions s	no wii odio w.	
	Less Frequently		Adverse
1072			Adverse
1072 1073	Less Frequently	Reported	
1072 1073 1074	Less Frequently Reactions	Reported events were	reported in
1072 1073 1074 1075	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney	Reported e events were ss than 15% in transplant reci	reported in acidence in pients who
1072 1073 1074 1075 1076	Less Frequently Reactions The following adverse the range of 3% to less	Reported e events were ss than 15% in transplant reci	reported in acidence in pients who
1072 1073 1074 1075 1076 1077	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney	Reported e events were ss than 15% in transplant reci	reported in acidence in pients who
1072 1073 1074 1075 1076 1077 1078	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac	Reported e events were ss than 15% in transplant reci	reported in acidence in pients who
1072 1073 1074 1075 1076 1077 1078 1079	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac comparative trials.	Reported e events were es than 15% in transplant reciprolimus in the SYSTEM:	reported in neidence in pients who e Phase 3
1072 1073 1074 1075 1076 1077 1078 1079 1080	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac comparative trials. NERVOUS	Reported e events were es than 15% in transplant reciprolimus in the SYSTEM:	reported in neidence in pients who e Phase 3
1072 1073 1074 1075 1076 1077 1078 1079 1080 1081	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac comparative trials. NERVOUS WARNINGS) abno amnesia, anxiety,	Reported e events were ss than 15% in transplant reciprolimus in the SYSTEM: srmal dreams	reported in neidence in pients who e Phase 3 (see , agitation, convulsion,
1072 1073 1074 1075 1076 1077 1078 1079 1080 1081 1082	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac comparative trials. NERVOUS WARNINGS) abno amnesia, anxiety, depression, dizzines	Reported e events were es than 15% in transplant reciperolimus in the SYSTEM: ermal dreams confusion,	reported in neidence in pients who e Phase 3 (see , agitation, convulsion,
1072 1073 1074 1075 1076 1077 1078 1079 1080 1081 1082 1083	Less Frequently Reactions The following adverse the range of 3% to les either liver or kidney were treated with tac comparative trials. NERVOUS WARNINGS) abno amnesia, anxiety, depression, dizzines encephalopathy, hal	Reported e events were ss than 15% in transplant recipations in the SYSTEM: symmal dreams confusion, s, emotional lucinations,	reported in neidence in pients who e Phase 3  (see , agitation, convulsion, l lability,

1087	abnormal; SPECIAL SENSES: abnormal vision,
1088	amblyopia, ear pain, otitis media, tinnitus;
1089	GASTROINTESTINAL: anorexia, cholangitis,
1090	cholestatic jaundice, dyspepsia, dysphagia,
1091	esophagitis, flatulence, gastritis, gastrointestinal
1092	hemorrhage, GGT increase, GI perforation,
1093	hepatitis, ileus, increased appetite, jaundice, liver
1094	damage, liver function test abnormal, oral
1095	moniliasis, rectal disorder, stomatitis;
1096	CARDIOVASCULAR: angina pectoris, chest
1097	pain, deep thrombophlebitis, abnormal ECG,
1098	hemorrhage, hypotension, postural hypotension,
1099	peripheral vascular disorder, phlebitis,
1100	tachycardia, thrombosis, vasodilatation;
1101	UROGENITAL: (see <b>WARNINGS</b> )
1102	albuminuria, cystitis, dysuria, hematuria,
1103	hydronephrosis, kidney failure, kidney tubular
1104	necrosis, nocturia, pyuria, toxic nephropathy,
1105	oliguria, urinary frequency, urinary incontinence,
1106	vaginitis; METABOLIC/NUTRITIONAL:
1107	acidosis, alkaline phosphatase increased, alkalosis,
1108	ALT (SGPT) increased, AST (SGOT) increased,
1109	bicarbonate decreased, bilirubinemia, BUN
1110	increased, dehydration, GGT increased, healing
1111	abnormal, hypercalcemia, hypercholesterolemia,
1112	hyperlipemia, hyperphosphatemia, hyperuricemia,
1113	hypervolemia, hypocalcemia, hypoglycemia,
1114	hyponatremia, hypophosphatemia,
1115	hypoproteinemia, lactic dehydrogenase

1116	increase, weight gain; ENDOCRINE: (see
1117	PRECAUTIONS) Cushing=s syndrome, diabetes
1118	mellitus; HEMIC/LYMPHATIC: coagulation
1119	disorder, ecchymosis, hypochromic anemia,
1120	leukocytosis, leukopenia, polycythemia,
1121	prothrombin decreased, serum iron decreased,
1122	thrombocytopenia; MISCELLANEOUS:
1123	abdomen enlarged, abscess, accidental injury,
1124	allergic reaction, cellulitis, chills, flu syndrome,
1125	generalized edema, hernia, peritonitis,
1126	photosensitivity reaction, sepsis;
1127	MUSCULOSKELETAL: arthralgia, cramps,
1128	generalized spasm, joint disorder, leg cramps,
1129	myalgia, myasthenia, osteoporosis;
1130	RESPIRATORY: asthma, bronchitis, cough
1131	increased, lung disorder, pneumothorax,
1132	pulmonary edema, pharyngitis, pneumonia,
1133	respiratory disorder, rhinitis, sinusitis, voice
1134	alteration; SKIN: acne, alopecia, exfoliative
1135	dermatitis, fungal dermatitis, herpes simplex,
1136	hirsutism, skin discoloration, skin disorder, skin
1137	ulcer, sweating.
1138	The overall safety profile of the Prograf-
1139	mycophenolate mofetil Phase IV study did not
1140	differ from the safety profile of the Phase III
1141	kidney study.

1143	
1144	Post Marketing
1145	The following have been reported: increased
1146	amylase including pancreatitis, hearing loss
1147	including deafness, leukoencephalopathy,
1148	thrombocytopenic purpura, hemolytic-uremic
1149	syndrome, acute renal failure, Stevens-Johnson
1150	syndrome, stomach ulcer, glycosuria, cardiac
1151	arrhythmia and gastroenteritis.
1152	There have been rare spontaneous reports
1153	of myocardial hypertrophy associated with
1154	clinically manifested ventricular dysfunction in
1155	patients receiving Prograf therapy (see
1156	PRECAUTIONS-Myocardial Hypertrophy).
1157	
1158	OVERDOSAGE:
1159	Limited overdosage experience is available. Acute
1160	overdosages of up to 30 times the intended dose
1161	have been reported. Almost all cases have been
1162	asymptomatic and all patients recovered with no
1163	sequelae. Occasionally, acute overdosage has
1164	been followed by adverse reactions consistent with
1165	those listed in the ADVERSE REACTIONS
1166	section except in one case where transient urticaria
1167	and lethargy were observed. Based on the poor
1168	aqueous solubility and extensive erythrocyte and
1169	plasma protein binding, it is anticipated that
1170	tacrolimus is not dialyzable to any significant
1171	extent; there is no experience with charcoal
1172	hemoperfusion.

1173	The oral use of activated charcoal has been
1174	reported in treating acute overdoses, but
1175	experience has not been sufficient to warran
1176	recommending its use. General supportive
1177	measures and treatment of specific symptoms
1178	should be followed in all cases of overdosage.
1179	In acute oral and IV toxicity studies
1180	mortalities were seen at or above the following
1181	doses: in adult rats, 52X the recommended human
1182	oral dose; in immature rats, 16X the
1183	recommended oral dose; and in adult rats, 16X
1184	the recommended human IV dose (all based or
1185	body surface area corrections).
1186	
1187	DOSAGE AND ADMINISTRATION:
1188	Prograf injection (tacrolimus injection)
1189	
1190	For IV Infusion Only
1191	
1192	NOTE: Anaphylactic reactions have
1193	occurred with injectables containing castor oi
1194	derivatives. See WARNINGS.
1195	
1196	In patients unable to take oral Prograf capsules
1197	therapy may be initiated with Prograf injection.
1198	The initial dose of Prograf should be administered
1199	no sooner than 6 hours after transplantation. The
1200	recommended starting dose of Prograf injection is
1201	0.03-0.05 mg/kg/day as a continuous IV infusion
1202	Adult patients should receive doses at the lower
1203	end

1204 of the dosing range. Concomitant adrenal 1205 corticosteroid therapy is recommended early post-1206 transplantation. Continuous IV infusion of Prograf 1207 injection should be continued only until the patient 1208 can tolerate oral administration of Prograf 1209 capsules. 1210 1211 1212 1213 Preparation for Administration/Stability 1214 Prograf injection must be diluted with 0.9% 1215 Sodium Chloride Injection or 5% Dextrose 1216 Injection to a concentration between 0.004 1217 mg/mL and 0.02 mg/mL prior to use. Diluted 1218 infusion solution should be stored in glass or 1219 polyethylene containers and should be discarded 1220 after 24 hours. The diluted infusion solution 1221 should not be stored in a PVC container due to 1222 decreased stability and the potential for extraction 1223 of phthalates. In situations where more dilute 1224 solutions are utilized (e.g., pediatric dosing, etc.), 1225 PVC-free tubing should likewise be used to 1226 minimize the potential for significant drug 1227 adsorption onto the tubing. Parenteral drug 1228 products should be inspected visually for 1229 particulate matter and discoloration prior to 1230 administration, whenever solution and container 1231 permit. Due to the chemical instability of 1232 tacrolimus in alkaline media, Prograf injection 1233 should not be mixed or co-infused with solutions 1234 of pH 9 or greater (e.g., ganciclovir or acyclovir).

1235
1236
1237 Prograf capsules (tacrolimus capsules)1238
1239 Summary of Initial Oral Dosage
1240 Recommendations and Typical Whole Blood

1241 Trough Concentrations

Patient Population	Recommended Initial Oral Dose*	Typical Whole Blood Trough Concentrations
Adult kidney transplant patients	0.2 mg/kg/day	month 1-3 : 7-20 ng/mL month 4-12 : 5-15 ng/mL
Adult liver transplant patients	0.10-0.15 mg/kg/day	month 1-12 : 5-20 ng/mL
Pediatric liver transplant patients	0.15-0.20 mg/kg/day	month 1-12 : 5-20 ng/mL

\*Note: two divided doses, q12h

1243 1244

#### Liver Transplantation

1245 It is recommended that patients initiate oral 1246 therapy with Prograf capsules if possible. If IV 1247 therapy is necessary, conversion from IV to oral 1248 Prograf is recommended as soon as oral therapy 1249 can be tolerated. This usually occurs within 2-3 1250 days. The initial dose of Prograf should be 1251 administered no sooner than 6 hours after 1252 In a patient receiving an IV transplantation. 1253 infusion, the first dose of oral therapy should be 1254 given 8-12 hours after discontinuing the IV 1255 infusion. The recommended starting oral dose of 1256 Prograf capsules is 0.10-0.15 mg/kg/day 1257 administered in two divided daily

grapefruit juice has been reported to increase

Co-administered

doses every 12 hours.

1258

1259

1260 tacrolimus blood trough concentrations in liver transplant patients. (See Drugs that May Alter 1261 1262 Tacrolimus Concentrations.) 1263 Dosing should be titrated based on 1264 clinical assessments of rejection and tolerability. 1265 Lower Prograf dosages may be sufficient as 1266 maintenance therapy. Adjunct therapy with 1267 adrenal corticosteroids is recommended early 1268 post transplant. 1269 Dosage and typical tacrolimus whole 1270 blood trough concentrations are shown in the 1271 table above; blood concentration details are 1272 described in **Blood Concentration Monitoring**: 1273 Liver Transplantation below. 1274 1275 Kidney Transplantation 1276 The recommended starting oral dose of Prograf 1277 is 0.2 mg/kg/day administered every 12 hours in 1278 two divided doses. The initial dose of Prograf 1279 may be administered within 24 hours of 1280 transplantation, but should be delayed until renal 1281 function has recovered (as indicated for example 1282 by a serum creatinine # 4 mg/dL). Black patients 1283 may require higher doses to achieve comparable 1284 blood concentrations. Dosage and typical 1285 tacrolimus whole blood trough concentrations are 1286 shown in the table above; blood concentration 1287 details are described in **Blood Concentration** 1288 Monitoring: Kidney Transplantation below.

The data in kidney transplant patients indicate that the Black patients required a higher dose to attain comparable trough concentrations compared to Caucasian patients.

Time After Transplant	Caucasian n=114		Black n=56	
	Dose (mg/kg)	Trough Concentration s (ng/mL)	Dose (mg/kg)	Trough Concentrations (ng/mL)
Day 7	0.18	12.0	0.23	10.9
Month 1	0.17	12.8	0.26	12.9
Month 6	0.14	11.8	0.24	11.5
Month 12	0.13	10.1	0.19	11.0

#### Pediatric Patients

Pediatric liver transplantation patients without pre-existing renal or hepatic dysfunction have required and tolerated higher doses than adults to achieve similar blood concentrations. Therefore, it is recommended that therapy be initiated in pediatric patients at a starting IV dose of 0.03-0.05 mg/kg/day and a starting oral dose of 0.15-0.20 mg/kg/day. Dose adjustments may be required. Experience in pediatric kidney transplantation patients is limited.

1307	
1308	
1309	Patients with Hepatic or Renal Dysfunction
1310	Due to the reduced clearance and prolonged half-
1311	life, patients with severe hepatic impairment (Pugh
1312	≥ 10) may require lower doses of Prograf. Close
1313	monitoring of blood concentrations is warranted.
1314	Due to the potential for nephrotoxicity, patients
1315	with renal or hepatic impairment should receive
1316	doses at the lowest value of the recommended IV
1317	and oral dosing ranges. Further reductions in
1318	dose below these ranges may be required.
1319	Prograf therapy usually should be delayed up to
1320	48 hours or longer in patients with post-operative
1321	oliguria.
1322	
1323	
1324	Conversion from One Immunosuppressive
1325	Regimen to Another
1326	Prograf should not be used simultaneously with
1327	cyclosporine. Prograf or cyclosporine should be
1328	discontinued at least 24 hours before initiating the
1329	other. In the presence of elevated Prograf or
1330	cyclosporine concentrations, dosing with the
1331	other drug usually should be further delayed.
1332	
1333	<b>Blood Concentration Monitoring</b>
1334	Monitoring of tacrolimus blood concentrations in
1335	conjunction with other laboratory and clinical
1336	parameters is considered an essential

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aid to patient management for the evaluation of rejection, toxicity, dose adjustments and compliance. Factors influencing frequency of monitoring include but are not limited to hepatic or renal dysfunction, the addition or discontinuation of potentially interacting drugs and the posttransplant time. Blood concentration monitoring is not a replacement for renal and liver function monitoring and tissue biopsies.

Two methods have been used for the assay of tacrolimus, a microparticle enzyme immunoassay (MEIA) and an ELISA. Both methods have the same monoclonal antibody for tacrolimus. Comparison of the concentrations in published literature to patient concentrations using the current assays must be made with detailed knowledge of the assay methods and biological matrices employed. Whole blood is the matrix of choice and specimens should be collected into tubes containing ethylene diamine tetraacetic acid (EDTA) anti-coagulant. Heparin anti-coagulation is not recommended because of the tendency to form clots on storage. Samples which are not analyzed immediately should be stored at room temperature or in a refrigerator and assayed within 7 days; if samples are to be kept longer they should be deep frozen at -20E C for up to 12 months.

1365 1366 1367 Liver Transplantation 1368 Although there is a lack of direct correlation 1369 between tacrolimus concentrations and drug 1370 efficacy, data from Phase II and III studies of 1371 liver transplant patients have shown an increasing 1372 incidence of adverse events with increasing trough 1373 blood concentrations. Most patients are stable 1374 when trough whole blood concentrations are 1375 maintained between 5 to 20 ng/mL. Long term 1376 posttransplant patients often are maintained at the 1377 low end of this target range. 1378 Data from the U.S. clinical trial show that 1379 tacrolimus whole blood concentrations, as 1380 measured by ELISA, were most variable during 1381 the first week post-transplantation. After this 1382 early period, the median trough blood 1383 concentrations, measured at intervals from the 1384 second week to one year post-transplantation, ranged from 9.8 ng/mL to 19.4 ng/mL. 1385 1386 Therapeutic Drug Monitoring, 1995, 1387 Volume 17, Number 6 contains a consensus 1388 document and several position papers regarding 1389 the therapeutic monitoring of tacrolimus from the 1390 1995 International Consensus Conference on 1391 Immunosuppressive Drugs. Refer to these 1392 manuscripts for further discussions of tacrolimus

1393

monitoring.

1394	
1395	
1396	Kidney Transplantation
1397	Data from the Phase III study indicates that
1398	trough concentrations of tacrolimus in whole
1399	blood, as measured by IMx7, were most variable
1400	during the first week of dosing. During the first
1401	three months, 80% of the patients maintained
1402	trough concentrations between 7-20 ng/mL, and
1403	then between 5-15 ng/mL, through one-year.
1404	The relative risk of toxicity is increased
1405	with higher trough concentrations. Therefore,
1406	monitoring of whole blood trough concentrations
1407	is recommended to assist in the clinical evaluation
1408	of toxicity.
1409	
1410	HOW SUPPLIED:
1411	Prograf capsules (tacrolimus capsules)
1412	0.5 mg
1413	Oblong, light yellow, branded with red "0.5 mg"
1414	on the capsule cap and " f 607" on the
1415	capsule body, supplied in 60-count bottles (NDC
1416	0469-0607-67) and 10 blister cards of 10
1417	capsules (NDC 0469-0607-10), containing the
1418	equivalent of 0.5 mg anhydrous tacrolimus.

1419	
1420	
1421	Prograf capsules (tacrolimus capsules)
1422	1 mg
1423	Oblong, white, branded with red "1 mg" on the
1424	capsule cap and " f 7" on the capsule
1425	body, supplied in 100-count bottles (NDC 0469-
1426	0617-71) and 10 blister cards of 10 capsules
1427	(NDC 0469-0617-10), containing the equivalent
1428	of 1 mg anhydrous tacrolimus.
1429	
1430	Prograf capsules (tacrolimus capsules)
1431	5 mg
1432	Oblong, grayish/red, branded with white "5 mg"
1433	on the capsule cap and " f 657" on the
1434	capsule body, supplied in 100-count bottles
1435	(NDC 0469-0657-71) and 10 blister cards of 10
1436	capsules (NDC 0469-0657-10), containing the
1437	equivalent of 5 mg anhydrous tacrolimus.
1438	
1439	Store and Dispense
1440	Store at 25°C (77°F); excursions permitted to
1441	15EC-30EC (59EF-86EF) [see USP Controlled
1442	Room Temperature].
1443	
1444	Prograf injection (tacrolimus injection) 5mg
1445	(for IV infusion only)
1446	Supplied as a sterile solution in 1 mL ampules
1447	containing the equivalent of 5 mg of anhydrous
1448	tacrolimus per mL, in boxes of 10 ampules (NDC
1449	0469-3016-01).

1450	
1451	
1452	Store and Dispense
1453	Store between 5EC and 25EC (41EF and 77EF).
1454	
1455	Rx only
1456	
1457	Made in Ireland
1458	for Fujisawa Healthcare, Inc.
1459	Deerfield, IL 60015-2548
1460	by Fujisawa Ireland, Ltd.
1461	Killorglin, Co. Kerry Ireland
1462	
1463	REFERENCE:
1464	1. CDC: Recommendations of the Advisory
1465	Committee on Immunization Practices: Use of
1466	vaccines and immune globulins in persons
1467	with altered immunocompetence. MMWR
1468	1993;42(RR-4):1-18.
1469	
1470	1/23/01
1471	
1472	Patient Information
1473	
1474	PROGRAF
1475	(tacrolimus capsules)
1476	
1477	
1478	Read this important information before you
1479	start using PROGRAF [PRO-graf] and
1480	each time you refill your prescription. This
1481	summary does not take the place of talking
1482	with your transplant team.
1483	
1484	Talk with your transplant team if you have
1485	any questions or want more information

1486	about PROGRAF. You can also visit the
1487	Fujisawa Internet site at www.fujisawa.com.
1488	
1489	What Is PROGRAF?
1490	
1491	PROGRAF is a medicine that slows down the
1492	body's immune system. For this reason, it
1493	works as an anti-rejection medicine.
1494	PROGRAF helps patients who have had a liver
1495	or kidney transplant protect their new organ
1496	and prevent it from being rejected by the body.
1497	
1498	How Does PROGRAF Protect My New
1499	Organ?
1500	
1501	The body's immune system protects the
1502	body against anything that it does not
1503	recognize as part of the body. For
1504	example, when the immune system detects
1505	a virus or bacteria it tries to get rid of it to
1506	prevent infection. When a person has a
1507	liver or kidney transplant, the immune
1508	system does not recognize the new organ
1509	as a part of the body and tries to get rid of
1510	it, too. This is called "rejection."
1511	PROGRAF protects your new organ by
1512	slowing down the body's immune system.
1513	
1514	Who Should Not Take PROGRAF?
1515	
1516	Do not take PROGRAF if you are allergic to
1517	any of the ingredients in PROGRAF. The
1518	active ingredient is tacrolimus. Ask your doctor
1519	or pharmacist about the inactive ingredients.
1520	-
1521	Tell your transplant team about all your health

conditions, including kidney and/or liver
problems. Discuss with your transplant team
the use of any other prescription and non-
prescription medications, including any herbal
or over-the-counter remedies that you may take
while on Prograf. In very rare cases you may
not be able to take Prograf.
Tell your transplant team if you are pregnant,
planning to have a baby or are breastfeeding.
Talk with your transplant doctor about possible
effects PROGRAF could have on your child.
Do not nurse a baby while taking PROGRAF
since the medicine will be in the breast milk.

1536		
1537		
1538	<b>How Shoul</b>	d I Take PROGRAF?
1539		
1540	PROGRAF	can protect your new kidney or
1541		you take the medicine correctly.
1542	•	•
1543	Your new o	rgan needs around-the-clock
1544		o your body does not reject it. The
1545	•	our transplant depends a great deal
1546	upon how w	vell you help PROGRAF do its job.
1547	_	t you can do to help.
1548		-
1549		
1550	\$	Take PROGRAF exactly as
1551		prescribed
1552		
1553		It is important to take
1554		PROGRAF capsules exactly as
1555		your transplant team tells you
1556		to.
1557		
1558		PROGRAF comes in several
1559		different strength capsules0.5
1560		mg, 1 mg and 5 mg. Your
1561		transplant team will tell you
1562		what dose to take and how
1563		often to take it. Your transplant
1564		team may adjust your dose until
1565		they find what works best for
1566		you.
1567		
1568		Never change your dose on
1569		your own. Never stop taking
1570		PROGRAF even if you are
1571		feeling well. However, if you

1572	feel poorly on Prograf, discuss
1573	this with your transplant team.
1574	· · · · · ·
1575	
1576	\$ Take PROGRAF two times
1577	a day, 12 hours apart
1578	
1579	Try to pick times that will be
1580	easy for you. For example, if
1581	you take your first dose at 7:00
1582	a.m. you should take your
1583	second dose at 7:00 p.m. Do
1584	not vary the times. You must
1585	take PROGRAF at the same
1586	times every day. If you decide
1587	to take PROGRAF at 7:00
1588	a.m. and 7:00 p.m., take it at
1589	these same times every day.
1590	This will make sure you always
1591	have enough medicine in your
1592	body to give your new organ
1593	the around-the-clock protection
1594	it needs.
1595	
1596	
1597	\$ Take PROGRAF the same
1598	way each day
1599	
1600	Some people prefer to take
1601	PROGRAF with food to help
1602	reduce possible stomach upset.
1603	Whether you take PROGRAF
1604	with or without food, it is
1605	important to take PROGRAF
1606	the same way every day. For
1607	example, if you take

1608 1609 1610 1611 1612 1613 1614 1615 1616 1617 1618 1619 1620 1621 1622	PROGRAF with food, you should always take it with food. Do not eat grapefruit or drink grapefruit juice in combination with your medicine unless your transplant teams approves. Do not change the way you take this medicine without telling your transplant team, since this could change the amount of protection you get from PROGRAF.
1623	\$ Take all your doses
1624	It is improved to tolk a vocal
1625	It is important to take your
1626	doses twice a day exactly as
1627 1628	prescribed by your doctor. If
1629	you miss even two doses, your
1630	new liver or kidney could lose the protection it needs to
1631	defend itself against rejection by
1632	your body.
1633	your body.
1634	If you miss one dose, do not try
1635	to catch up on your own. Call
1636	your transplant team right away
1637	for instructions on what to do.
1638	
1639	If you travel and change time
1640	zones, be sure to ask your
1641	transplant team how to adjust
1642	your dosage schedule so your
1643	new organ does not lose its

1644	protection.
1645	
1646	
1647	<ul> <li>Plan ahead so that you do</li> </ul>
1648	not run out of PROGRAF
1649	
1650	Make sure you have your
1651	prescription for PROGRAF
1652	refilled and at home before you
1653	need it. Circle the date on a
1654	calendar when you need to
1655	order your refill. Allow extra
1656	time if you receive your
1657	medicines through the mail.
1658	
1659	Your transplant team will follow your progress
1660	and watch for early signs of side effects. This is
1661	why you will have blood tests done often after
1662	your transplant. On the days you are going to
1663	have a blood test to measure the amount of
1664	PROGRAF in your body, your transplant team
1665	may ask you not to take your morning dose
1666	until after the blood sample is taken. Check
1667	with your transplant team before skipping this
1668	dose.
1669	
1670	
1671	Can Other Medicines Affect How
1672	PROGRAF Works?
1673	
1674	Some medicines and alcohol can affect how
1675	well PROGRAF works. After you start taking
1676	PROGRAF:
1677	
1678	<b>\$</b> Be sure to tell your transplant
1679	team, family doctor, dentist,

1680		pharmacist and any other health
1681		care professional treating you
1682		the names of <b>all</b> the medicines
1683		you are taking. This includes
1684		PROGRAF as well as all other
1685		prescription medicines and non-
1686		prescription medicines, natural
1687		or herbal remedies, nutritional
1688		supplements, and vitamins. This
1689		is the only way that your health
1690		care team can help prevent
1691		drug interactions that could be
1692		serious.
1693		
1694	\$	Always check with your
1695		transplant team before you start
1696		taking any new medicine.
1697		
1698	\$	While you are taking
1699		PROGRAF, do not get any
1700		vaccinations without your
1701		transplant team's approval.
1702		The vaccination may not work
1703		as well as it should.
1704		
1705	\$	Liver transplant patients,
1706		including those taking
1707		PROGRAF, should not drink
1708		alcohol.
1709		
1710	What Are the	Possible Side Effects of
1711	PROGRAF?	
1712		
1713		
1714		
1715	Tell your transp	plant team right away if you think

1716	you might be having a side effect. Your
1717	transplant team will decide if it is a medicine
1718	side effect or a sign that has nothing to do with
1719	the medicine but needs to be treated. Infection
1720	or reduced urine can be signs of serious
1721	problems that you should discuss with your
1722	transplant team.
1723	
1724	Your transplant team will also follow your
1725	progress and watch for the early signs of any
1726	side effects. This is why you will have blood
1727	tests done often during the first few months after
1728	your transplant. On the days you are going to
1729	have a blood test to measure the amount of
1730	PROGRAF in your body, your transplant team
1731	may ask you not to take your morning dose
1732	until after the blood sample is taken. Check
1733	with your transplant team before skipping this
1734	dose.
1735	
1736	
1737	
1738	For Kidney Transplant Patients:
1739	
1740	The most common side effects of
1741	PROGRAF for kidney transplant
1742	patients are infection, headache,
1743	tremors (shaking of the body), diarrhea,
1744	constipation, nausea, high blood
1745	pressure, changes in the amount of
1746	urine, and trouble sleeping.
1747	
1748	Less common side effects are
1749	abdominal pain (stomach pain),
1750	numbness or tingling in your hands or
1751	feet; loss of appetite; indigestion or

1752	"upset stomach"; vomiting; urinary tract
1753	infections; fever; pain; swelling of the
1754	hands, ankles or legs; shortness of
1755	breath or trouble breathing; cough; leg
1756	cramps; heart "fluttering", palpitations
1757	or chest pain; unusual weakness or
1758	tiredness; dizziness; confusion; changes
1759	in mood or emotions; itchy skin, skin
1760	rash, and diabetes.
1761	
1762	
1763	For Liver Transplant Patients:
1764	
1765	The most common side effects of
1766	PROGRAF for liver transplant patients
1767	are headache, tremors (shaking of the
1768	body), diarrhea, high blood pressure,
1769	nausea and changes in the amount of
1770	urine.
1771	
1772	Less common side effects are
1773	numbness or tingling in your hands or
1774	feet; trouble sleeping; constipation; loss
1775	of appetite; vomiting; urinary tract
1776	infections; fever; pain (especially in the
1777	back or abdomen [stomach area]);
1778	swelling of the hands, ankles, legs or
1779	abdomen; shortness of breath or
1780	trouble breathing; cough; unusual
1781	bruising; leg cramps; heart "fluttering"
1782	or palpitations; unusual weakness or
1783	tiredness; confusion; changes in mood
1784	or emotions; itchy skin, and skin rash.
1785	
1786	

Be sure to tell your transplant team right

1787

1788	away if you notice that you are thirstier
1789	than usual, have to urinate more often,
1790	have blurred vision or seem to get
1791	confused. These may be the early signs of
1792	high blood sugar or diabetes.
1793	
1794	All anti-rejection medicines, including
1795	PROGRAF, suppress your body's immune
1796	system. As a result, they may increase your
1797	chances of getting infections and some kinds of
1798	cancer, including skin and lymph gland cancer
1799	(lymphoma). As usual for patients with
1800	increased risk for skin cancer, exposure to
1801	sunlight and UV light should be limited by
1802	wearing protective clothing and using a
1803	sunscreen with a high sun protection factor
1804	(SPF \$ 15). However, getting cancer from
1805	taking an anti-rejection medicine is not
1806	common. Talk with your transplant team about
1807	any concerns or questions you have.
1808	
1809	
1810	How Should I Store PROGRAF?
1811	
1812	Store PROGRAF in a dry area at room
1813	temperature (77° F/25° C). Do not let the
1814	medicine get colder than 59° F (15° C) or
1815	hotter than 86°F (30°C). For instance, do not
1816	leave PROGRAF in the glove compartment of
1817	your car in the summer or winter. Do not keep
1818	PROGRAF capsules in a hot or moist place
1210	such as the medicine cabinet in the bathroom

1820	
1821	
1822	
1823	
1824	
1825	General Advice about Prescription
1826	Medicines
1827	
1828	Medicines are sometimes prescribed for
1829	conditions that are not mentioned in patient
1830	information leaflets. Do not use PROGRAF for
1831	a condition for which it was not prescribed. Do
1832	not give PROGRAF to other people.
1833	
1834	This leaflet summarizes the most important
1835	information about PROGRAF. If you would
1836	like more information, talk with your doctor.
1837	You can ask your pharmacist or doctor for
1838	information about PROGRAF that is written for
1839	health professionals. You can also visit the
1840	Fujisawa Internet site at www.fujisawa.com.
1841	
1842	
1843	Fujisawa logotype
1844	[address, copyright, date, code, etc.]
1845	
1846	